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1 Development and evolution of tooth renewal in neoselachian
2 sharks as a model for transformation in chondrichthyan
3 dentitions

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17 Key words: Chondrichthyan, teeth, dentitions, replacement, evolution

18
19 **Short title: *Neoselachian transformations from alternate***
20 ***replacement tooth order***

21
22 **Abstract**

23 A defining feature of dentitions in modern sharks and rays is the
24 regulated pattern order that generates multiple replacement teeth.
25 These are arranged in labio-lingual files of replacement teeth, that form
26 in sequential time order both along the jaw, and within successively
27 initiated teeth in a deep dental lamina. Two distinct adult dentitions
28 have been described; one as alternate, in which timing of new teeth
29 alternates between two adjacent files, each erupting separately. The
30 second is different, arranged as single files, where teeth of each file
31 are timed to erupt together, in some taxa facilitating similarly-timed
32 teeth to join to form a cutting blade. Both are dependent on
33 spatiotemporally regulated formation of new teeth.

The adult Angel shark *Squatina* (Squalomorphii) exemplifies a single file dentition, but we obtained new data on the developmental order of teeth in the files of *Squatina* embryos, showing alternate timing of tooth initiation. This was based on micro-CT scans revealing that the earliest mineralised teeth at the jaw margin and their replacements in file pairs (odd and even jaw positions) alternate in their initiation timing..

Along with *Squatina*, new observations from other squalomorphs such as *Hexanchus* and *Chlamydoselachus*, together with representatives of the sister group Galeomorphii, have established that the alternate tooth pattern (initiation time and replacement order) characterises the embryonic dentition of extant sharks, but this can change in adults. These character states were plotted onto a recent phylogeny, demonstrating that the Squalomorphii shows considerable plasticity of dental development. We propose a developmental-evolutionary model to allow change from the alternate to a single file alignment of replacement teeth. This establishes new dental morphologies in adult sharks from inherited alternate order.

Introduction

In modern-grade elasmobranch fishes, Neoselachii (including all living sharks, rays and skates), replacement tooth positions along the jaw have been described as two distinct arrangements, single file or alternate file (e.g. Reif 1980; Smith et al. 2013; Underwood et al. 2015; Fig. 1A, C), or modifications of these basic patterns, to produce for example, a single cutting blade of imbricated teeth across the jaw (Strasburg 1963; Underwood et al. 2015). During development, an asynchronous timed series of tooth germs is initiated along the jaw (Smith et al. 2009), at alternate positions, as labio-lingual files of replacement teeth (Figs. 1D, 2). In this arrangement of replacement teeth, two adjacent files form (Smith 2009; Smith et al. 2013, fig. 1, 2), from the first rudimentary teeth, as sequentially added teeth (SAT unit), proposed as a clonal unit of differentially timed teeth (SAT unit

of tf 6+7, Fig. 2). In these, timing and position alternate within each pair of tooth files (tf 6+7) to provide closely spaced teeth in alternately timed labio-lingual addition (Fig. 1B). Each pair of files are added disto-proximally along the jaw (Fig. 2, 1-12), developmentally linked as iteratively timed 'SAT units' (Fig. 2, stages 1-9; Smith 2003; Smith et al. 2013: figs 1A, 2), so that in the adult teeth erupt at different times. By comparison, the single file arrangement in the adult is composed of teeth in adjacent files at the same developmental phase, recognised by the youngest teeth being at the same stage of development and the oldest at the same position relative to the jaw margin, erupting together (Fig. 1A). In this arrangement, timing of eruption at the jaw margin could be either as single teeth or, as simultaneous for all teeth forming a blade (Fig. 1A; Underwood et al. 2016), and as alternate teeth (Fig. 1B). Dentitions may also have disto-proximally staggered times (Fig. 2, 1-12), and many rows of erupted (functional) teeth together (Fig. 1C). Until recently, it was unclear how these different adult morphologies developed and if neoselachian dentitions showed developmental plasticity, through transformation of tooth order from embryo to adult, and whether alternate or single file addition is the primitive state for skates and rays

Within the Batoidea (skates and rays), the sister group to modern sharks (Squalomorphii + Galeomorphii, Fig. 3), both embryos and adults possess the alternate pattern for arrangement of successor teeth. Within many batoids teeth in alternating order are close packed, forming continuous surfaces, for crushing dentitions (Underwood et al. 2015). In sharks, all members of the Galeomorphii so far studied (Carcharhiniformes, Orectolobiformes and Heterodontiformes; see Smith et al. 2013; Fig. 1B, D) have alternate tooth replacement in adults, from alternate developmental order in embryos. The status of the Lamniformes has been considered as uncertain (Smith et al. 2013). In Squalomorphii, many taxa have distinctive single-file successor teeth demonstrated to be the result of developmental modification of an embryonic alternate pattern (Underwood et al. 2016; Fig. 1A). The

alternate pattern can also be present in the upper jaw, or both jaws, as an example of developmental independence (plasticity) of upper and lower jaws, as in Hexanchidae (lower dentition single file, but alternate pattern in the upper jaw and teeth that lie closest to the jaw hinges).

Several other clades of squalomorph sharks appear to have single file tooth replacement order in adults, with well-spaced tooth files, including *Squatina* (angel shark), the Hexanchidae (six and seven gilled sharks), and *Chlamydoselachus* (frilled shark), as well as the Lamniformes within the Galeomorphii (Mako, Thresher and White sharks and relatives), but in each case their developmental order is unknown. Our study of the embryonic and adult dentitions of *Squatina*, along with examination of dentitions of *Hexanchus*, *Chlamydoselachus* and *Isurus* (Galeomorphii), allows the early stages of tooth development to be compared, and contrasted with the embryonic alternate patterning in *Heterodontus* (Heterodontiformes), as detailed by Reif (1976).

These data are used to explore the hypothesis that the alternate pattern can be transformed into single-file during development both through alteration of tooth germ timing and loss of individual tooth files, with these being two different potential mechanisms for evolutionary transformation. We propose that the alternate addition of teeth is plesiomorphic for the Neoselachii, retained in embryonic and adult rays, and present in fossil relatives, but can be modified to single file from the embryonic condition in some adult sharks (Galeomorphii and Squalomorphii). Embryonic *Squatina* is an excellent model to test this hypothesis in squalomorphs along with *Chlamydoselachus* and *Hexanchus* as basal members of the group (Fig. 3) all with putatively single file dentitions in the adult.

To determine the plesiomorphic state, fossils of taxa closely related to extant sharks and rays were studied, including *Acrodus* (Hybodontoidae) and *Synechodus*. The phylogenetic relationships of *Synechodus* are uncertain (Maisey 1985; Klug 2010; Fig. 3, where 'Neoselachii' includes *Synechodus* + Elasmobranchii), although it shares many characters with extant sharks and rays, including that of a complex tooth histology (e.g., Enault et al. 2015). These two taxa

were investigated with the aim of testing whether the alternate arrangement of teeth in adults is the basal state for modern sharks as well as rays (Fig. 3), and where clades with the derived state (single file replacement) occurred on the phylogeny.

Materials and methods

Materials

Wet specimens

Squatina californica, embryo (Natural History Museum, Life Sciences collection, BMNH 91.5.19.240)

Isurus oxyrinchus, embryo (BMNH 1961.11.2.3)

Chlamydoselachus anguineus, adult jaw (BMNH 2016.4.11.1)

Lamna nasus, adult head, jaw sections (BMNH 2015.3.13 1-3)

Hexanchus sp. (?*H. nakamurai*), embryo (BMNH 1973.7.12.4-6)

Fossil specimens

Synechodus dubrisiensis, jaw (Booth Museum, Brighton, BMB 008523)

Acrodus anningae, jaw (Natural History Museum, Earth Sciences collection, NHMUK PV P2732)

Dried specimens (Birkbeck reference collection)

Squatina spp. (*Squatina guggenheim*, *S. argentina*, *S. tergocellata*, *S. ?caillaiti*)

Echinorhinus brucus

Hexanchidae (4 species; *Hexanchus griseus*, *H. nakamurai*, *Heptranchias perlo*, *Notorynchus cepedianus*)

Lamniformes (8 species; *Lamna nasus*, *Isurus oxyrinchus*, *Pseudocarcharias kamoharai*, *Carcharias taurus*, *Odontaspic ferox*, *Alopias pelagicus*, *A. superciliosus*, *A. vulpinus*.)

Cleared and stained specimens

Chlamydoselachus anguineus embryos and juvenile (Nos. 1984/5/6/6, 1984/9/2/3, and 1985/5/3C: TL 28.6cm; 40.1 cms). These were stained with Alizarin red, Alcian blue, from the Tokai University Museum, Shimizu, Japan (TMFE), courtesy of Sho Tanaka, also studied (Nos. 1984/5/6/6, 1984/9/2/3, and 1985/5/3C: TL 28.6cm; 40.1

168 cms).

169 **Methods**

170 *Imaging*

171 We used x-ray computed tomography to examine the head region of
172 whole embryos (micro-CT, Nikon Metrology HMX ST 225, Image and
173 Analysis Centre, Natural History Museum, London (NHM)) to visualise
174 the teeth present within the jaws of specimens, especially the earliest
175 teeth (mineralised cusps) from the 3D volume rendered models.
176 Photomacrographs were taken using a Nikon Coolpix camera in natural
177 light; drawings made with the software ‘Inkscape’ and the X11 Window
178 System.

179 *Terminology*

180 For use of directional terms such as distal and proximal, see
181 Underwood et al. (2016). The systematic terminology follows
182 Compagno (1973, 1977) and Nelson (2006).

183 **Measurements**

184 In embryonic *Squatina*, measurements were taken when the first
185 initiated teeth are set iteratively along the jaw (Fig. 4, 1-8, distal to
186 proximal) with 3-4 labio-lingual successive teeth and none shed from
187 the jaw margin (Figs. 4, 5, stage 1, 2, 3). We compared this data to
188 embryos of *Chlamydoselachus* and *Isurus* (taxa also with apparent
189 single file tooth replacement). We also investigated various adult
190 galeomorph and squalomorph taxa, along with fossil dentitions,
191 selected from key positions on a neoselachian phylogeny (Fig. 3).

192 Tooth size was measured in two different ways, depending on the
193 overall morphology of the tooth. The first focused on cusp height, as
194 the distance between the tip of the central cusp to the crown/base
195 boundary. Distances between the selected points were calculated using
196 Avizo 9.2 software's measurement tools. A 3D rendered image was
197 used for *Squatina* (embryo and adult) and *Chlamydoselachus* (adult)
198 with the distance between the two points being recorded for each
199 tooth in the file, starting at the first file next to the symphysis. Thus
200 each tooth file includes separate measurements of tooth cusp height
201 starting with the oldest tooth (labial) to the youngest developing

mineralized tooth (lingual).

The second method, where tooth file size differed along the jaw, as in the *Isurus* embryo, and the central cusp was asymmetric relative to the base, used a different approach. Here, tooth volume was estimated to overcome problems such as the wide range of asymmetrical variation of each tooth file (Fig. SI 4). Measuring volume required individually segmented tooth elements (Avizo 9.2), performed slice by slice, selecting mineralized regions of each tooth. A label analysis tool was then used to acquire an individual tooth volume metric ($\times \text{mm}^3$).

Histograms (Figs. 5G-I, 6H; Figs. SI 4, 6) produced used a colour code indicate developmental order for each tooth; darker colours represent the oldest tooth, grading to the lightest colours for the developmentally youngest teeth. Also, within each tooth file, red and green represent odd and even tooth files respectively, beginning at the symphyseal region.

Results

Developmental interpretation of tooth replacement

In chondrichthyans, the developmental dental lamina restricts tooth induction (odontogenic potential) to the tissues in the lamina (Reif 1982; Smith et al. 2009, and references therein; Rasch et al. 2016; Martin et al. 2016), which is present along the lingual face of the jaw cartilage. Here, tooth files are established in the embryo from the initiator teeth ordered sequentially along the jaw margins, where morphogenesis gradually assumes the adult shape, and in the adult new teeth are continually initiated (Figs. 1B, D, 2, pg). Because of this, developmental interpretations can be made from observations of static morphology, with teeth effectively suspended in a time sequence (Reif 1980, 1982). Hence the smallest, oldest teeth are at the jaw margin, while the newest teeth are in various stages of development deep in the lingual furrow (Figs. 1D, 2). Newest or youngest teeth are visible only as the mineralized central cusp tip, but in each tooth file they form in a sequentially timed developmental

order with increasing morphological differentiation (e.g., cusps joined to base), to become the largest in the tooth file. These observations from tooth germs in the dental lamina can be used to assign relative timed order to sequential stages of the developing teeth within two adjacent files (Figs. 1D, 2, arrows).

Squalomorphii

***Squatina* embryonic dentition**

In embryos, as in adults, teeth were arranged in well-spaced single files, with 10 files in the upper jaw and 11 in the lower, in each half (Figs. 4, 5, Fig. SI 2C). In the embryonic lower jaw, the youngest developing teeth occur lingually, as small cusps (no.8, Figs. 4, 5B, D, E, asterisks), with the oldest teeth labially, and towards the jaw symphysis. These have a single central cusp joined to a small tooth base (nos. 1&2 in Figs. 4, 5D). A single rudimentary symphyseal tooth is present in one specimen (red, Figs. 4, 5E, F), but was not observed in others. This is only present in the first tooth row, with subsequent rows lacking symphyseal teeth. When all tooth files along the jaw are established (disto-proximally, Fig. 4), it is apparent that the oldest (first) teeth are the smallest and nearest to the labial jaw margin, and located in even positions relative to the symphyseal tooth (Fig. 5, t1, t2, B, D). These teeth are about to erupt, or be shed (purple, Fig. 4; Fig. 5A, C, alternate pink). Thus, teeth in even number positions indicate the first initiation time in each pair of alternate files and of first loss from the jaw margin. Similarly, those at odd positions (Fig. 4, green) are the second teeth to be initiated; then alternate sequential tooth initiation within the pairs (odd and even files, Fig. 2) has occurred with the latest to form, the newest tooth germ, visible as a cusp tip (Figs. 4, 5D no. 8, 5B, E, asterix).

The first four teeth of the four files were compared, with the fourth still a developing tooth (Fig. 5G, I). When represented as graphs comparing cusp volumes in rows parallel to the jaw margin, even number first teeth are smaller up to row three (green, Fig. 5H). In

row 1, as noted, the first teeth are smallest in even number files (2, 4) representing the first to form. In row 4, all developing teeth have only the central cusp, in even number files (green) cusps are larger than odd, because these are the first file teeth to develop and are now further advanced morphologically at this stage of development (row 4, Fig. 5H).

All teeth are central cusp aligned (Fig. 5C, short pink line), in progressive states of morphological development of central cusp and the tooth base in files numbers t1-7, t2-8 (Fig. 4). Each file contains four teeth, with two in the most proximal files (presumed newest files added proximally to the jaw, Fig. 4).

Interpretation of developmental timing in embryonic tooth files - As described above, teeth in files 1 and 2 on the right lower jaw illustrate relative sizes and morphologies, and therefore developmental order of timing for sequential tooth addition, alternating between these files (Fig. 5B, D-F). The first tooth of the series is the most labial and smallest formed tooth with an attachment base; sequential addition starts from file 2 (Fig. 5B, t1-7), tooth number 1 being formed before the morphogenetic program was perfected (Fig. 5B, D, E). The sequential time series ends with the newly initiated tooth tip of tooth number 8, as a mineralised but incomplete central cusp (Fig. 5B, E, asterix). This alternate file developmental set represents the SAT unit (Fig. 2, sat. tf 6+7). A single file is equivalent to one even number set (Fig. 2, sat. tf2). Order of initiation (t1-8, Figs. 4, 5D) may also determine timing of shedding, as the labial positions of first teeth in each alternate file indicate an alternate shedding order (pink, Fig. 5A, C). In each separate file central cusps are aligned (Fig. 5C, short pink line), also only alternate tooth cusps align along a straight disto-proximal line along the jaw, showing that adjacent files are offset (Fig. 5C, long pink line).

Measurements of central cusp heights and alignment - Morphological evidence of progressive, developmental tooth order as

clonal SAT units (paired odd and even files) was tested with measurements of tooth size, both along the jaw in rows parallel to the jaw margin, and within files (Fig. 5C, pink lines). These are represented as histograms (Fig. 5H, I) taken from tooth files 1-4 (alternate red and green, Fig. 5F, G vignettes of two alternate files). The comparative central cusp heights, represented by the graphs, confirm the smallest, complete first teeth are in the even number files (green, Fig. 5G, I), while the first tooth in odd number files is larger (red, Fig. 5H, row 1). When represented as graphs comparing cusp heights in rows parallel to the jaw margin, again even number first teeth are smaller up to row three (green, Fig. 5H); in row 4 teeth the central cusp is just developing and even numbers are larger than odd, because as these are the first of the teeth to develop they are further advanced in morphology at this stage of development (row 4, Fig. 5H).

Adult jaws of *Squatina guggenheim*

In the early adult dentition, occlusion of upper jaws with lowers shows teeth fitting between lowers, with teeth organized in widely spaced files along the jaw (Fig. SI 2B-D), so that tooth file spacing allows upper teeth to fit between lower ones. From the relative cusp heights it appears that the alternate initiation of teeth seen in an embryo does not continue in the adult (Fig. SI 2F); measurements in the adult dentition were taken, as for the embryonic teeth, from the first teeth of each of four files; these showed little difference between alternate positions (the fifth and sixth teeth are smaller (partially developed) so cannot be compared with the four fully formed main cusp volumes). However, in proximal files closer to the jaw articulation (Fig. SI 2C, white box, D, file numbers 6-10), the position of the oldest teeth located at the jaw margins seems to alternate and suggests they still show different times of origination, as in the embryonic alternate developmental pattern.

Other Squalomorphii

***Chlamydoselachus anguineus* embryo and adult dentitions**

In the embryo, at the 4-tooth stage (Fig. 6A-D) teeth are superficial and encapsulated in individual sheaths of connective tissue, separate for each file (Fig. 6A, contrast enhanced density). A symphyseal file is fully expressed in the lower jaw (Fig. 6A, B), but absent in the upper jaw (Fig. 6C). In the dried specimen (Fig. 6B), the symphyseal file has a rudimentary single cusp tooth that is nearest of all files to the jaw margin, and as such is the initiator tooth of the lower jaw dentition, to either side is also a single cusp first tooth, each file then acquires morphological competence as three cusped teeth. This morphology (Fig. 6C-G) is demonstrated with calcium positive staining (red, Fig. 6C-E) first as separate cusps (Fig. 6D, arrow, 6C), later joined to the base. In lower jaw files three and four (Fig. 6D), the smallest first tooth (oldest, asterix) is positioned in the even number file four, whereas, the third file first tooth is slightly larger (i.e. began developing later). The third and fourth teeth of file 4 have developing, mineralized cusps (4th tooth largest, as latest and most morphogenetic competence). These size differences in adjacent files suggest the presence of alternate tooth initiation timing. In the upper jaw at the four-tooth stage in each file, teeth are absent from the symphysis, the first tooth of each has only one cusp and the base has formed. Measurements of tooth sizes were not possible in these embryos, as they were only soft tissue preparations, or were long term fixed specimens and had lost mineral.

The lower jaw of a more mature embryo has tooth files proximally that had not completed morphogenesis (Fig. 6E), as all files have a rudimentary first labial tooth, so the embryo has not shed the first teeth in these proximal files and had not reached maturity (see Fig. 6F). These gradually reduce in total tooth number in files to 4 or 3, and all diminishing in size (Fig. 6E). In all files, tooth six is the youngest, with only cusp tips mineralized, but teeth in older positions have all cusps joined and the first four teeth also have cusps joined to mineralized bases. In the adult lower jaw these proximal files still have minute teeth but the first tooth is three-cusped (Fig. 6E), on the labial side of the jaw and was not rudimentary. We segmented and measured

the teeth of these last four files (Fig. 6F) to test whether we could show size differences that represented ordered, alternate files created by alternate timing, as in the embryo. The general tooth sizes of the four measured files (1-4 in Fig. 6F, H) decreases proximally, nevertheless, the first teeth were smaller in green files relative to the red files (Fig. 6F, H). We interpret this to result from the alternate developmental program seen in the embryo, still present in adult proximal teeth, but this needs to be tested on more suitable material.

Considering the fully formed teeth of the adult lower jaw, distal to these proximal files (Figs. 6G, Fig. SI 3), each file has similarly-sized teeth with four teeth fully erupted and locked together as a functional unit, lingually a fifth tooth has an incomplete base whilst a sixth tooth is forming as three separate cusps lingual to the completed teeth (Fig. 6G). These developing successional teeth are located on a lingual shelf on Meckel's cartilage deep to the oral surface (not in a separate bulla), older teeth are held in the connective tissue of the skin, as in the virtual section next to one of the tooth files (Fig. SI 3A). In the symphyseal file and three files either side, tooth size differences between odd and even number files were inconspicuous (Fig. SI 3C, histogram). It would seem that any evidence for timing difference between tooth files seen in the embryos is not present in the adult dentition, but may be present in the smallest, most proximal tooth files.

Hexanchidae: *Hexanchus* spp. embryo and adult dentitions; *Notorhynchus cepedianus*, adult dentitions

A single embryo of *Hexanchus* was studied, in which the teeth are well developed, but with those in the first row of the lower jaw only starting to rotate into a pre-functional position (Fig. 7A, C). Thus, most teeth have not been lost, as bulk shedding only occurs after the replacement teeth are in their functional positions (Underwood et al. 2016). The lower teeth are large and distinct, but no obvious alternation or overlapping of tooth bases can be observed, nor in the upper jaw (Fig. 7B).

In the embryo, individual replacement teeth in seven adjacent single files, including the symphyseal file, are obliquely arranged within disto-proximal rows (Fig. 7C, red lines). Although, tooth size decreases to the most proximal, seventh tooth file, equivalent individual teeth in files are approximately equal in size. This oblique, developmental alignment of teeth in a disto-proximal row has also been recognized in the Squalomorphii, demonstrated to be as a result of realignment of teeth to form a single file arrangement, altered from the embryonic alternate arrangement of the first teeth (Underwood et al. 2016). By comparison, in *Hexanchus*, embryonic teeth align as single file and no detectable alternation of replacement teeth is apparent. This arrangement is retained in the adult, where teeth are also single file aligned labio-lingually (e.g., Smith et al. 2013: fig. 4G).

An abrupt change in tooth form and arrangement occurs in the lower dentition of adult hexanchid genera, where the most proximal teeth are very reduced in size, and appear to show an irregular but alternating pattern (*Notorhynchus*, Fig. 7D). These reduced teeth in the adult are suggested to be a ‘remnant’ of the ancestral developmental alternate pattern (as described above for *Chlamydoselachus*), but not observed in *Hexanchus*, where alternate tooth addition was absent even in the embryo. This suggests that dental arrangement in the hexanchids is highly plastic, which appears to be a general feature of Squalomorphii, as discussed below (Fig. 3).

Galeomorphii

As the sister group to the Squalomorphii, we also considered the distribution of alternate file versus single file tooth addition for Galeomorphii, focusing on the Lamniformes (Fig. 3) and one embryo of Heterodontiformes, but excluding Orectolobiformes and Carcharhiniformes, both of which have alternate tooth replacement, as previously described (see Smith et al. 2013).

***Isurus oxyrinchus*, embryo (Lamniformes)**

The dentition in the embryo of *Isurus* comprises teeth of adult-like

morphology (Fig. 8A-E) in the oldest and smallest fully mineralized teeth but none have rotated into a functional position (except two teeth, see below). Tooth files 1 to 3 are held within a small, distal bulla next to the jaw symphysis, and more proximal files within a longer bulla, the two being separated by a space, or diastema (between files 3 and 4, Fig. 8C, E). In the adult of *Lamna*, (Fig. 8F) the disto-proximal number of tooth files (1-13) is the same as in the embryo (Fig. 8C-E, 1-13), indicating that the latter is a fully formed, unerupted dentition.

Despite tooth size varying dramatically along the jaw, size measurements of the first 5 files either side of the jaw symphysis (taken as in *Squatina*) did not show differences between teeth in odd versus even files (Fig. SI 4). However, this analysis did indicate the presence of a developmentally missing file (number 3 on each side). As mentioned above, alternate timing of tooth development in adjacent files can also be assessed via the relative position of the oldest teeth in each file relative to the jaw margin, and the overlap of root base lobes. This was far less problematic in the upper jaws than lower, in part due to the more oval cross section of the Meckel's cartilage making assessment of tooth proximity to the jaw margin less certain, hence assessment was made on the upper dentition.

In the upper dentition, the tooth in file 2 is oldest (Fig. 8E, red) as the only representative of the most labial disto-proximal tooth row, but without other teeth; evidence of shedding is shown by a tooth of identical morphology that has become lodged in the branchial region (Fig. 8A, white circle). The second row of alternating teeth includes teeth in files 1, 5, 7, 10 and 12 (Fig. 8C-E, yellow). The third tooth row includes teeth in files 2, 3, 4, 6, 8, 9, 11 and 13. This pattern is irregular (i.e. teeth in files 3 and 4 and in files 8 and 9 do not alternate in position), and as such forms a partial single file dentition, because both alternating tooth replacement and some regions of single file tooth replacement occur at this stage of development. The most likely explanation for the highly specific regional lack of alternating files (corroborating the graphic data, Fig. SI 4A, C) is that files have been developmentally lost. For example, the diastema between distal and

more proximal teeth could mark the position of one of the missing tooth files.

***Lamna nasus*, adult (Lamniformes)**

In the upper dentition in adults of *Lamna*, tooth files are also held in proximal and distal bullae, with an intervening diastema (between 3 and 4) even more prominent than in the *Isurus* embryo. In the upper jaw, the oldest teeth in files 3 and 4 are at the same position relative to the jaw margin (Fig. 8F). Whilst the relative positions of the distal tooth files are not clear, teeth in files 8, and 9, do not appear to alternate and likewise have their oldest teeth at the same position relative to the jaw margin and their youngest teeth at the same stage of development. The teeth within the upper jaw of an adult *Lamna* thus resemble the *Isurus* embryo, alternate file differential in timing, but small portions of the dentition showing single file replacement due to regional, probable loss of intervening tooth files (e.g., at the diastema).

***Alopias pelagicus*, adult (Lamniformes)**

The upper dentition of adult *Alopias pelagicus* has the lamniform arrangement of three tooth files positioned within an anterior bulla, although this bulla is smaller and less distinct than in *Isurus* and *Lamna*. In *Alopias*, tooth files show an alternate pattern of eruption times, or tooth bases, relative to the jaw margin (Fig. 8G, H, files 1 and 3, and 2 and 4). Also, the oldest teeth in files 5 and 7 (Fig. 8H, 4 and 6, and 5 and 7) are in the same position relative to the jaw margin, as evidence of alternate timing events.

***Heterodontus* sp. (Heterodontiformes)**

In a late stage embryo (labial view, Fig. 9A), teeth have rolled over the jaw margin, with the symphyseal tooth and one tooth present from file 2 in the initial row, with these being the earliest to form. Here arrangement is alternate (Fig. 9A) but file two on the right jaw may have shed a tooth, consistent with tooth shedding in the earliest

embryos, as described previously (Reif 1976: fig. 8F). Teeth with cusped morphology in distal files have a bilateral symmetry (as in symphyseal teeth) and alternate in their position with respect to the jaw margin, as do the more proximal molariform teeth with broad bases and low cusps (tooth positions 7-9), in the less curved part of the jaw (Fig. 9B-D). New teeth are added proximally to molariform teeth at jaw position 10, as a large open base to the developing but narrow crown (Fig. 9E, arrows). In the adult (Smith et al. 2013: fig. 3A, B), the close packing of both tooth morphologies seems to present as 'single file' but in the embryo the molariform teeth are slightly staggered in each file, to accommodate wide teeth formed in an alternate pattern.

Fossil taxa

***Synechodus* (Synechodontiformes)**

The fossil shark genus *Synechodus* is generally considered to be part of a monophyletic clade of neoselachians (Synechodontiformes; Klug 2010), although this is not universally accepted (e.g. Maisey 1985). Whilst most specimens of *Synechodus* and other Synechodontiformes comprise isolated teeth, several skulls and well-preserved skeletons are known (Maisey 1985). Within these, however, dentitions are commonly poorly exposed or partly disarticulated. In contrast, a dentition from the Late Cretaceous of SE England (Fig. 10A-I) comprises an articulated suite of teeth with no jaw cartilages preserved. This shows a high degree of heterodonty (examples as isolated teeth, Fig. 10J-L) with erect and cusped teeth distally (Fig. 10D, G-I) and wide low cusped teeth in more proximal positions (Fig. 10A-C, E, F, L), but all are arranged with bases and crowns in close packed, alternate arrangement. At the symphysis (Fig. 10D, H) two files of relatively small, close packed parasymphyseal teeth clearly alternate in their positions, labio-lingually. This alternate packing of *in situ* teeth in all positions of this fossil specimen indicates that the dentition derives from an alternating tooth addition pattern at their initiation.

***Acrodus* (Hybodontoidae)**

The extinct Hybodontoidae are a clade forming a sister group to Neoselachii (Maisey 1987; Lane 2010; Fig. 3). Observations were made on an exceptionally well-preserved dentition of *Acrodus* from the Early Jurassic of southern England (Fig. 10M, N). This specimen preserves an entire dentition but, as with the *Synechodus* specimen, no jaw cartilages are preserved. CT-renders revealed some additional teeth (unprepared from the fossil matrix) on the labial margin of the dentition and in the process of shedding. It also showed the presence of pyrite, an iron sulphide that forms early in the fossilization process, between the teeth, suggesting that teeth have not shifted relative to each other *postmortem*. The exposed biting surface of the dentition reveals wide, low teeth on either side of a file of smaller teeth at the jaw symphysis. The extended tooth bases in adjacent files overlap, demonstrating an alternate pattern of replacement teeth. Observations of other hybodont sharks such as *Asteracanthus* (Rees and Underwood 2006) show that a similar alternating pattern was present across the clade.

Discussion

The aims of our study were focused on the spatio-temporal order of initiation of replacement teeth in the Neoselachii. The relative timing of successive teeth in the files was determined from observations and measurements of relative size differences of teeth, compared both within and between adjacent tooth files, in adults and embryos wherever possible.

Focusing on embryos of the Angel shark *Squatina* (SI Figs. 1, 2), we have proposed a model for all neoselachians (i.e. iterative sequence of tooth addition), from a structural pattern of the dentition that we interpret as timed developmental differences based on a clonal generative unit (SAT unit) that comprises two adjacent tooth files (Fig. 2). This developmental model with alternate timing and sequential spatial arrangement explains the development of the alternate tooth pattern (Smith 2003; Smith et al. 2013). It is

proposed that the alternate pattern of tooth addition (based on Reif's observations (1978): Fig. 2) is plesiomorphic for the Neoselachii (Fig. 3); as a phylogenetically basal neoselachian developmental pattern that operates in the first stages of the embryonic dentition, it results in alternately timed eruption and shedding. We examined specific taxa in the Squalomorphii and Galeomorphii, to see how this model may be altered during development to arrange teeth as a single file tooth pattern in adults, including simultaneous eruption into a cutting blade (Underwood et al., 2016).

In *Squatina* (Squalomorphii), we have demonstrated that the earliest dentition shows alternate timing (clonal SAT units, Figs. 4, 5) and although tooth size differences are not apparent in the adult (suggesting a shift to a single file pattern), an alternate arrangement appears to be retained in the young adult indicated by their positions at the proximal jaw margins (Fig. SI 2C, D). Although well-separated single files are present in the adult of *Chlamydoselachus*, similarly rudimentary proximal tooth files in the adult retain an alternate arrangement (Fig. 6F, H), but we have only scarce data to show that embryos exhibit alternate tooth addition, in their early development. A better understanding of how tooth addition timing changed from embryo to adult in these taxa requires gene expression data, including how co-ordination and alignment of tooth files between upper and lower jaws is programmed.

The Hexanchidae, closely related to *Chlamydoselachus* (Fig. 3), have a unique and highly distinctive dentition, with tooth morphology very different from that of *Chlamydoselachus* and *Squatina*. The lower teeth of adult Hexanchidae are arranged in a single file order, as in the embryo of *Hexanchus*, but each tooth is wide and abuts the next so together they form a 'saw blade' at the jaw margin (see Smith and Johanson 2015: fig. 1.7), as in other Squaliformes (Underwood et al. 2016). But unlike the Squaliformes and *Squatina* (Fig. 3), this arrangement does not appear to derive from an alternate arrangement in the embryo in *Hexanchus*. However, in hexanchid adults such as *Notorhynchus*, the most proximal rudimentary teeth do retain a

developmental alternate arrangement (Fig. 7D: as in the adults of *Squatina* and *Chlamydoselachus*), as evidence of an underlying, persistent inherited alternate tooth order.

The mode of tooth replacement in the Lamniformes, a very specialized group of sharks including both macropredators and planktivores with diverse dentitions reflecting their differing trophic roles, has been uncertain (SI movies; Smith et al. 2013) due to the widely spaced tooth files and high degree of curvature of the jaw cartilages towards the symphysis. They are unique in that the dentition of some species is functional long before birth, and used during intrauterine oophagy and cannibalism, so that earliest stages of the tooth order are uncertain (e.g. Shimada 2002; Tomita et al. 2017). In predatory lamniforms the upper dentition is distinctive (Fig. 8F), with tooth files originating in deep proximal and distal bulla, the latter comprising a suite of typically three files. Between these is a raised cartilage bar, which developmentally may relate to a diastema formed by the loss of tooth files (in the Lamniformes), or have one or more files of very reduced teeth (*Mitsukurina*, *Carcharias*), varying between individuals or between the jaws of the same individual.

We noted that the development of teeth within lamniforms is a two-stage process, and only the second phase of tooth growth is addressed here. Embryos develop an early suite of teeth (Shimada 2002; Tomita et al. 2017) that is shed prior to the eruption of the adult-type teeth (Fig. 8A, white circle). Although the tooth arrangement in the first dental set is as yet unclear (in part due to the rarity of these embryos) we have shown that the tooth size differs in adjacent files and therefore has an alternate timing of tooth addition (Figs. 8, SI 4). Overall, *Isurus* and other Lamniformes show an alternate pattern that is modified in early development through loss of tooth files, near the diastema in the upper jaw and elsewhere, resulting in portions of the jaw possessing single file tooth replacement (Fig. 3, state 2).

Phylogenetic relationships

Recent molecular phylogenies of extant sharks support the sister group relationship between the major clades Squalomorphii and Galeomorphii, with these forming a sister group to the Batoidea (e.g., Velez-Zuazo and Agnarsson 2011; Fig. 3). Phylogenetically, *Synechodus* has been resolved as a sister taxon to extant sharks and rays (Klug et al. 2009; Klug 2010), but also assigned to the Galeomorphii (Maisey 2012; Fig. 3). Although teeth of *Synechodus* are organized in close packed files (central cusp labio-lingually aligned), the tooth bases overlap, reflecting alternate tooth initiation and eruption. An alternate replacement was also demonstrated for the hybodont *Acrodus* (Hybodontoidae, also resolved as being a sister taxon to extant sharks and rays; Fig. 3). Clearly defined alternate tooth replacement patterns are also present in embryos and adults of both the Galeomorphii (Smith et al. 2013) and Squalomorphii (e.g. Pristiophoridae; Underwood et al. 2016). Along with this, dentitions in the batoids show alternate tooth addition, both within the embryo and adult (Underwood et al. 2015; Fig. 3). These observations suggest that the alternate pattern of tooth addition is plesiomorphic for the Neoselachii as a whole.

Single file tooth replacement in certain dentitions of Squaliformes has been recognized as a derived modification of this alternate pattern during development (Underwood et al. 2016; Fig. 3, state 3). *Squatina* and *Chlamydoselachus* show separate tooth files in the adult but retain an alternate pattern of tooth addition, at least proximally (Fig. 3, state 3). Separation of tooth files may have resulted from a fixed number of tooth files (new files not added proximally) combined with jaw growth, or from tooth file loss, nevertheless these taxa show a markedly different mechanism than that suggested in the Squaliformes (Underwood et al. 2016).

By comparison, in the Hexanchidae, single file tooth replacement is present in both the embryo and adult of *Hexanchus*, although remnants of an alternate dentition are preserved in files of reduced teeth in adult hexanchids like *Notorhynchus*. *Isurus* and other Lamniformes show a different developmental pattern for replacement,

involving modification through loss of whole tooth files near the diastema, resulting in single file tooth replacement limited to this region (e.g., previously alternate files at the same developmental stage are now adjacent to one another).

These modified patterns are uncommon in the neoselachians, representing a derived state acquired from the embryonic alternate dentition, but by different mechanisms; in the Lamniformes by loss of tooth files (Fig. 3, state 2) and within squalomorph clades by independent modification of the alternate differential timing along the jaw (Fig. 3, state 3; see also Strasburg (1963) for dentition modification from an alternate tooth arrangement). These taxa appear to have developmental plasticity sufficient to allow the formation of single file tooth replacement by re-ordering of tooth production from an embryonic pattern of sequential, alternate timing of tooth initiation. Heterodontiformes, as the most basal taxon in the Galeomorphii, exhibit distinct alternate arrangement in the embryo, and retention in the adult (Fig. 9).

Conclusions

This study has investigated the initial development of tooth replacement patterns in a number of shark taxa, where successional tooth order was previously poorly understood. We speculate that a change in timing of replacement tooth addition, or loss of tooth files, resulted in the shift from embryonic to adult dentitions and loss of the alternate pattern in some taxa. These changes allowed teeth to emerge simultaneously at the jaw margin, forming a continuous cutting edge as an adaptation to a specific feeding mode, or functionally driven adaptation.

In addition, we studied tooth replacement series in adults of two extinct species of sharks, representing sister taxa to extant groups. We mapped characters onto a recent phylogeny based on transformation, or retention, of a developmental process (interpreted for fossil species) into ‘single file’ or ‘alternate file’ ordering of replacement teeth in the adult. In this way we predict the basal

(alternate) and derived (single file) phylogenetic states and suggest how this evolved through ‘tinkering’ with developmental mechanisms but by differing methods in the Squalomorphii, and the Galeomorphii.

Thus, the combination of fossil and extant phylogenetic data suggests that the alternate tooth pattern is plesiomorphic for the Neoselachii, with modification in adult sharks, but achieved differently in the two major clades. Squalomorphs modified an embryonic alternate tooth replacement pattern in various ways to generate a single file pattern in the adult, but by loss of tooth files in the galeomorphs. We postulate that these groups have the developmental plasticity to allow the formation of single file tooth replacement via the re-ordering of tooth production. This was not a substantial mechanism within the Neoselachii, which otherwise were dominated by alternate patterning of the dentition.

Hence, we propose a developmental-evolutionary model from the alternate pattern to achieve a single file alignment of replacement teeth, one with coequal eruption times at the functional edge of the jaws.

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746 Author Contributions

747 MMS conceived this study, MMS, BC, ZJ, CU generated micro-CT scan
748 data, JK provided data on *Chlamydoselachus*, all authors contributed to
749 the interpretation of data, and the writing of the paper.

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751 Conflict of Interests

752 We declare that we have no conflicts of interest.

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FIGURE LEGENDS

Fig. 1. Adult and embryonic jaws with tooth arrangement at the jaw margin

Single file (A) versus alternate dentitions (B, C), with (D) unknown timed order in embryo. (A) *Scymnodon ringens* (Knifetooth dogfish), lower jaw in labial view with single, symmetrical tooth across the jaw symphysis (S). (B) *Prionace glauca* (Blue shark), upper jaw in lingual view with replacement row teeth including symmetrical symphyseal tooth, other teeth are polarised left-right (modified from Smith et al., 2013, figs. 1B, 5C; photos Tom Diekwisch). (C) *Triaenodon obesus* (White tip reef shark) adult dentition with alternate dentition, reduced symphyseal teeth. (D) *Carcharhinus leucas* (Blacktip shark) embryonic lower jaw (lingual epithelium removed, tissues stained) with tooth files central cusp aligned but all successor teeth appear in single file arrangement from rudimentary cusp of first tooth (volumetric data not available) with space for attachment bases to increase in size (from Smith, 2003: fig. 9A; Smith et al. 2013: fig 4D). Scale bars= 1.0cm.

Fig. 2. Developmental model of dentition in alternate file order in Grey reef sharks Carcharhinidae, single cusp teeth are first initiated along the jaw, formed as mineralized tissue in embryos with one tooth row (stage1), then two rows (stage 2), later in development nine tooth rows (stage 9). Jaw positions (distal to proximal) numbered 1-12 from the symphyseal tooth (S), first as even number positions, then odd in the second row. Smallest teeth (black, stage 1) then larger alternate teeth with polarised shape (grey, stage 2), later larger teeth with lateral cusps form by row 3 (Smith et al., 2013, fig. 2). Sequential tooth initiation in a clonal set (arrows, direction of timing for teeth, 1-9) shows the alternate timing of tooth initiation order in adjacent tooth files 6 & 7 (SAT unit tf 6+7), with the next putative tooth germ (pg) to form in odd number row position. An example as if it was a single file sequential addition model is superimposed on this alternate model at file position 2 (sat tf 2; Smith et al. 2013: fig. 2; modified from Reif, 1978).

Fig. 3. Neoselachian phylogeny with character state distribution

Alternate and single tooth file replacement in embryos and adults, with these character states plotted on a recent phylogeny (Naylor et al. 2012). The basal position and monophyly of the Synechodontiformes, including *Synechodus*, follows Klug (2010). All neoselachians, as well as the Hybodontoidae, show alternate tooth replacement in some part of their dentition in the embryo, even if this is not retained in the adult (state 1). Within the Lamniformes (Galeomorphii) some tooth files are lost to produce the appearance of single file tooth addition, found in only certain parts of the jaw, reflecting irregular tooth file loss (state 2). Within the Squalomorphii, the single file tooth replacement pattern is developed from secondary modification of an alternate pattern (state 3). Within *Chlamydoselachus* an alternate pattern may be present in the embryo but not in the adult (except proximally); the majority of the dentition shows a single file arrangement. In *Hexanchus* (Hexanchidae), single file addition is present in early development and retained in the adult, while in other adult hexanchids (e.g., *Notorhynchus*) the alternate pattern is only retained in proximal rudimentary teeth. In *Squatina* an alternate pattern is present in the embryo, while the adult dentition possesses a single file arrangement but has retained alternate addition in proximal tooth files. In these three taxa state 3 (single file) may be related to a fixed number of tooth files and independent jaw growth, and in *Squatina* allows interdigitation of upper with lower jaw teeth.

Fig. 4. Schematic drawing of timed developmental order of first dentition teeth

Left lower jaw of *Squatina californica* embryo (BMNH91.5.19.240), teeth in apparent single file arrangement (from xCT render, Fig. 5) with odd (green) and even number (purple) tooth files and symphyseal tooth (red); timing order of clonal units shows successional tooth initiation (1-8) between first two adjacent files (based on size and morphology of files t1-7, and t2-8 in, Fig. 5A-E). Teeth at the labial jaw margins are rudimentary, first to form on the jaw cartilage at the even numbered jaw positions and closer to eruption and shedding. Newest lingual tooth (8) is smallest in odd numbers, representing mineralised central cusp tip. Arrows indicate labio-lingual tooth addition and disto-proximal tooth file addition.

Fig. 5. Spaced tooth files in lower jaw of *Squatina californica* embryo

BMNH 91.5.19.240, comparison of micro-CT 3D renders using VGStudio Max (B, D, E) and Avizo (A, C, F). (A) Labial view of right jaw, alternate position of rudimentary teeth at the jaw margin (pink), nearest are even number positions, white box indicates field in B, D. Labial view from symphysis (B, E) and lingual view (D), indicate developmental series of replacement teeth in time series from 1-8 (D). (B) First file (t2-8) and second (t1-7), at even tooth positions (t1-7) is first rudimentary tooth, at odd position (t2-8) are youngest (asterix, 8). (C) Oblique anterior view shows left jaw with alternate positions of shedding teeth (pink), right jaw central cusp aligned file (short pink line), alternate file cusps aligned (long pink line). (D) Tooth files 1, 2 as in B, E with sequence of initiation time order t1-8, tooth tip 8 the latest to form (see fig. 4). (E) Symphyseal region, labial view of tooth files in B, D, symphyseal tooth at jaw margin (red), newest tooth in replacement series of file 1 (asterix, 8 in D). (F) First four files segmented as alternate odd (red), even positions (green), from symphyseal tooth (sy t), volume of central cusps measured. (G) Colour profiles of first four teeth shown in each file in histogram, darkest is first in file (I). (H) Histogram showing relative sizes of individual teeth in rows (labio-lingual) first row smallest. (I) Histogram shows relative heights of each tooth in each alternate file, first is smallest in even numbers (colours as in (G)). Scale bars =5mm (B, C, F).

Fig. 6. Single file dentition of embryo and adult of

Chlamydoselachus angineus

(A) Micro-CT scan through dissected symphyseal segment of the embryo lower jaw dentition, tissue contrast enhanced with phosphotungstic acid. Symphyseal file with three tooth files either side, each with 5 tooth germs, separately encapsulated in connective tissue, the last has three developing cusps as in adult teeth (G). (B) Two views, lower jaw region across the symphysis of alcohol dried, younger embryo, isosurface render of three tooth files, symphyseal with smallest, single cusp, nearest of the three files to labial margin as first tooth formed, tooth files left and right also have single cusp first, but larger than the first symphyseal, second teeth (in all three) are larger single cusps and two small lateral cusps. (C D, E) Photomicrographs of cleared and stained embryos (Alcian Blue, Alizarin Red); (C) Upper jaw symphysis lacking symphyseal tooth file, first file (left) has small rudimentary first tooth of one cusp, second with 2 cusps, third with 3 cusps, with base outlined, fourth with 3 large cusps not joined at base; (D) Files 2 and 3 of lower jaw may show smallest first tooth in even number files (asterix), and in same file the fourth tooth has developing central cusps (arrow), larger than in adjacent file. (E) Juvenile, proximal eight files, reducing tooth numbers proximally, first teeth in all files rudimentary, increasing overall size distally. (F, G) Rendered and segmented adult lower jaw (BMNH2016.4.11.1) (F) Lingual view, smallest, but most proximal ordered tooth files (1-4 used for volume measurements, (H) Contrast seen with small cluster of oropharyngeal denticles lacking organisation and demal denticles, top, (see (G), and Fig. SI 3B). (G) Lingual view, four more distal files in which all teeth are above the jaw cartilage (no separate bullae), only loosely held in connective tissue (see Fig. SI 3A). (H) histograms of most proximal tooth files (coloured inset; files 1-4 in (F)); labial tooth is darkest colour in each set), insignificant size differences seen between first tooth in green files relative to red. Scale bars (A)= 3 mm.

Fig. 7. Single file dentition of embryo of *Hexanchus*

(A-C) *Hexanchus ?nakamurai* (BMNH1973.7.12.4-6), micro-CT renders of late stage embryo, upper and lower jaws. (A) Lower jaw dentition labial view, compared with (B) upper jaw dentition. (C) Lower right, lingual view (of A), three teeth in each file (file 7 has 2), teeth are aligned in a single file replacement pattern, but within each file they are arranged at an oblique

angle relative to the jaw margin (red line). (D) Proximal teeth of adult *Notorynchus cepedianus*, adult lower dentition, adjacent to last tooth of typical bladeliike morphology, tiny rudimentary teeth showing and alternate arrangement. Scale bar A, B = 5mm, C = 2.5mm. D, Scale bar= 10mm.

Fig. 8. Single file embryonic dentitions of *Isurus*, adult *Lamna* and *Alopias* (Lamniformes)

(A-E) *Isurus oxyrinchus* late stage embryo (BMNH1961.11.2.3), 3D-rendered micro-CT images. (A) Braincase, jaws, anterior vertebral column, lateral view. Developing teeth are visible in single file organisation and a loose tooth (upper right, white circle) in the gill region demonstrates that teeth are being shed at this stage. (B) Upper jaw region, internal view, partially erupted teeth, red. Tooth loss is confirmed by comparison between the fully erupted second tooth (black arrow), with a gap in the corresponding position on the right (black asterisk). (C) Left upper dentition, oldest teeth in adjacent files are at different positions relative to the jaw margin (see colour scheme, (E)). (D) Upper jaw, tooth rows in oblique lateral view, note the youngest teeth are in alternate even numbered files. (E) Upper left dentition with the oldest teeth colour coded to show their relative timing of development (see above). The tooth in file 2 is most developed (red) and probably oldest (relates to adjacent odd number files being the younger of each alternate pair, as in alternate model). Yellow teeth represent the next oldest with blue teeth being younger. Note the lack of alternation between files 3-4 (diastema) and 8-9 as presumed missing files. (F) *Lamna nasus* adult, upper dentition, diastema between 3-4. Teeth in most adjacent files alternate, but this is not seen in files 8-9. (G, H) *Alopias pelagicus*, macrophotos of adult, upper dentition, (G) Right upper dentition showing the typical lamniform arrangement of three teeth within an anterior bulla. (H) Part of the left upper jaw of the same individual; extra tooth file present in position 3. Scale bars = 10 mm.

Fig 9. *Heterodontus* sp. late stage embryo, 3D-rendered micro-CT images. (A) Lower dentition, labial view, file 2 of the right jaw shows a lost tooth (black arrow) from same row as symphyseal tooth (S). (B) Lingual view. (C) Left jaw, lingual view showing high-cusped distal teeth (jaw positions 1-6) and molariform teeth (jaw positions 1-9)), all in alternate arrangement. (D) Closeup of lower jaw, lingual view, tooth files showing

alternate pattern, both sides of symphyseal tooth file. (E) Lower jaw, oblique lingual view, latest tooth position (10) with tooth germs added proximally (white arrows). Scale bars, A, C-E= 5mm, B= 1cm.

Fig. 10. Articulated dentition of *Synechodus dubrisiensis* (Cretaceous, Chalk)

(A-I, BSB008523, Booth Museum, Brighton) A, E, H, macrophotos, B-D, F, G, I, 3D rendered micro-CT images. (A) Occlusal view, teeth are articulated as *in situ* but no cartilages are preserved. (B) Rock surrounding tooth files removed virtually. (C) Oblique lingual view showing the general arrangement of the teeth. (D) Detail of the symphyseal region showing alternating small teeth. (E) Lingual view, wide, low crown shape and alternating arrangement. (F) Same region, with pseudocoloured crowns to show close packed alternating pattern of adjacent files. (G) Teeth with high cusps (as in H-K) close to the symphysis show an alternating pattern of adjacent files contra those more proximally. (H) Symphyseal region with alternation of parasymphyseal teeth and large teeth on adjacent files. (I) Oblique view of the dentition, showing that several tooth rows would have been simultaneously functional. Scale bars, A-C, I = 10mm, D-H = 5mm. (J-L) *Synechodus dubrisiensis* isolated teeth (Early Cretaceous, Underwood 1999: pl. 1.1, 1.4, 1.5), SEM images, demonstrating the high degree of heterodonty. (J) Symphyseal tooth, (K) High cusped anterior tooth, (L) Low crowned posterior tooth. Scale bars = 1mm. (M, N) *Acrodus anningae* (NHMUK PV P2732), articulated dentition, as prepared (M) compared with segmented micro-CT image with rock removed virtually showing new teeth lingually (N). Pyrite is indicated in yellow. Note the alternating pattern of the tooth bases.

Supplementary Information

Fig. SI 1 Single file dentition in occlusion in late stage embryo of *Squatina californica*

(A-D) 3D rendered micro-CT images. (A) Occluded jaws, anterior view. (B) Meckel's cartilages and lower teeth (green), occlusal view, overlain by palatoquadrate and upper teeth (red), showing how upper and lower tooth files alternate in position along the jaw in occlusion. (C) Right palatoquadrate and upper teeth (blue), occlusal view, overlain by Meckel's cartilage and lower

teeth (red). (D) Right palatoquadrate and upper teeth (blue), and Meckel's cartilage and lower teeth (red), lateral view. Scale bars, A= 1cm, B= 3cm.

Fig. SI 2 Single file dentition of *Squatina guggenheim* small adult.

(A-D) 3D-rendered micro-CT images. (A) Dried skull, labial cartilages removed, dorsal view of lower jaw view, Scale bar = 50mm. (B) Left upper dentition with alternate tooth files red and green, central cusp aligned, lingual view. Scale bar = 10mm. (C, D) Lower dentition, (D) from field in box (C), proximal teeth show alternate positions at the jaw margin (white arrows). (E) Symphyseal region of the lower tooth files 1-4, odd (red), even (green), dorsal view. (F) Histograms of relative cusp heights of the four tooth files on either side of the symphysis, (as in (E), darker colours oldest tooth). There is little significant height differences between first teeth of the odd and even files, although file 4 is overall smaller.

Fig. SI 3 Single file dentition of *Chlamydoselachus anguineus*, adult

(A-B) 3D-rendered micro-CT images. (A) Lower jaw, developing tooth files on Meckel's cartilage in slight lingual furrow where tooth germs are forming, with virtual section of a tooth file in profile, functional teeth rolling over the jaw margin, held in soft tissue. (B) Oblique lateral view of Meckel's cartilages, teeth as spaced single files, up to five functional at the same time, skin denticles very small labially. Scale bars, A, B =10 mm. (C) Histograms of three files left and right side of symphysis (blue), as depicted in inset, as sizes of 5-7 teeth in each file, darkest shade represents first teeth in file; these being of even size.

Fig. SI 4 Dentition of lower jaw of *Isurus* embryo.

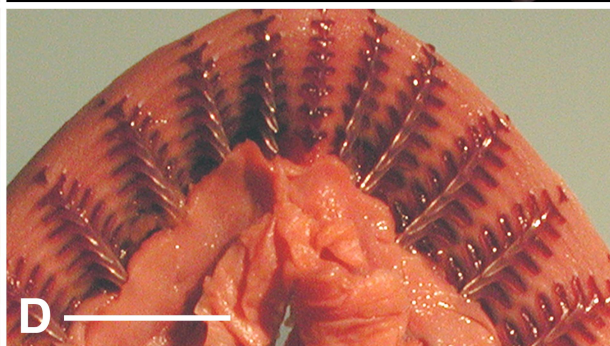
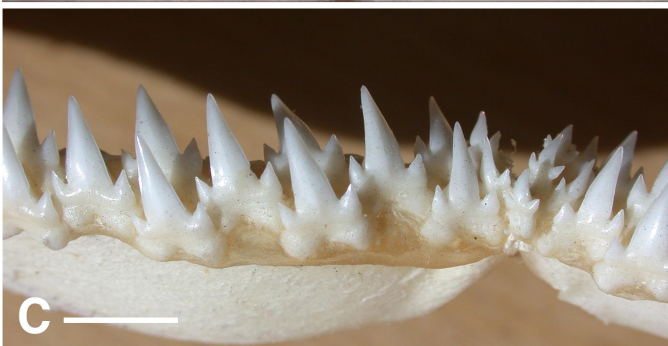
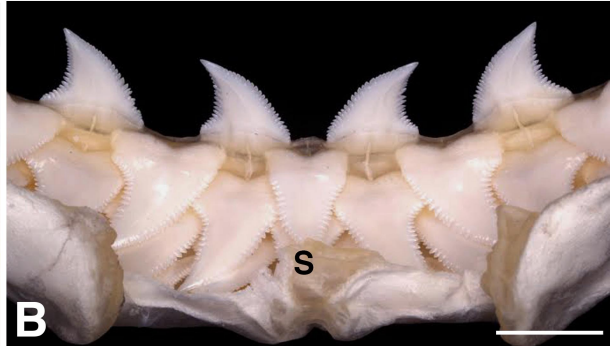
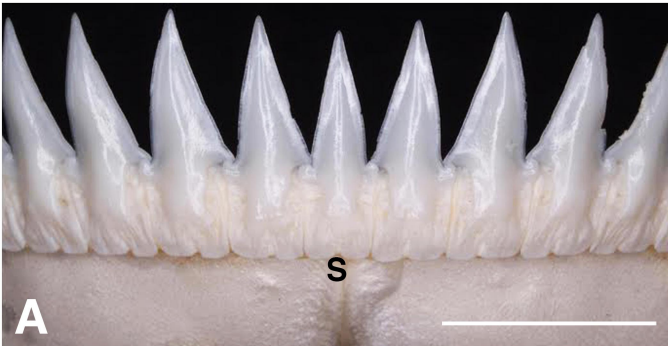
(BMNH1961.11.2.3) (A) Histogram showing relative tooth volume (mm^3) of each stage tooth in each alternate file. First five files depicted as alternate odd (red), even positions (green); darker shades representing the oldest, lightest shades the youngest. (B) Lower jaw, lingual view (Micro-CT 3D render, Avizo). Tooth files showing relative volume of the first five files, using eight equal bin sizes starting with the smallest, $0-0.444 \text{ mm}^3$ (dark blue) to the largest, $3.115-3.56 \text{ mm}^3$ (white). (C) Line graph showing relative tooth volume (mm^3) of each tooth row; Row 1 being the oldest and Row 4 with newly forming developing teeth. Scale bar =10mm (B).

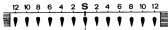
1087 **Fig. SI 5 Adult *Lamna nasus* (BMNH2015.3.13 1-3), movie generated**
1088 **from 3D rendered micro-CT scan.** Partial dissection of the tissues away
1089 from the whole dentition demonstrates alignment of teeth at the jaw margin
1090 and all files of replacement teeth.

1091

1092 **Fig SI 6 Adult *Lamna nasus* (BMNH2015.3.13 1-3), movie generated**
1093 **from 3D rendered micro-CT scan, showing teeth organized in a whorl.**

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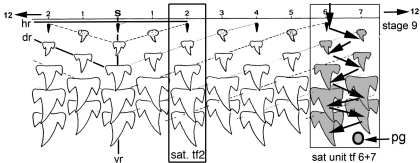




stage 1



stage 2

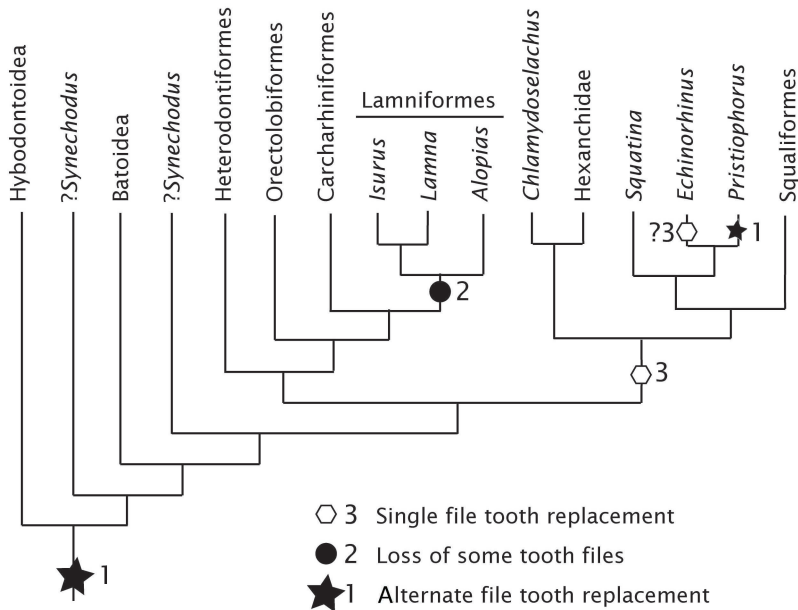


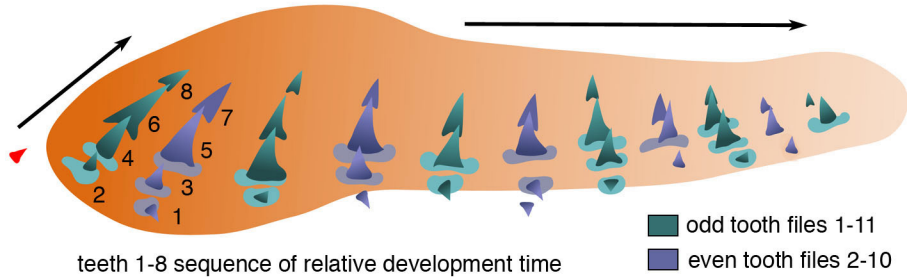
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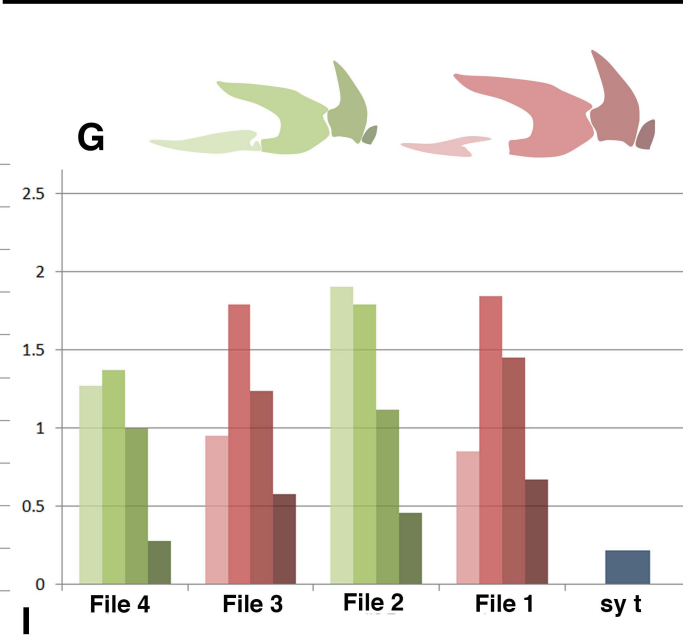
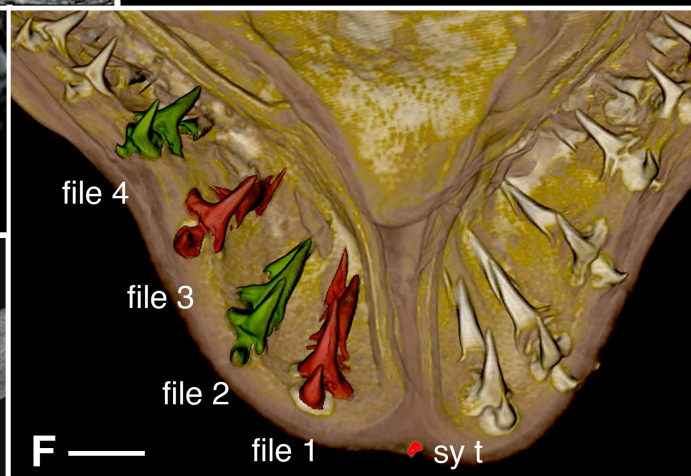
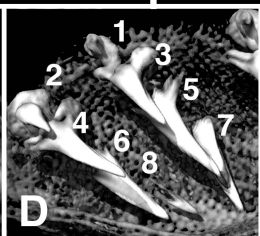
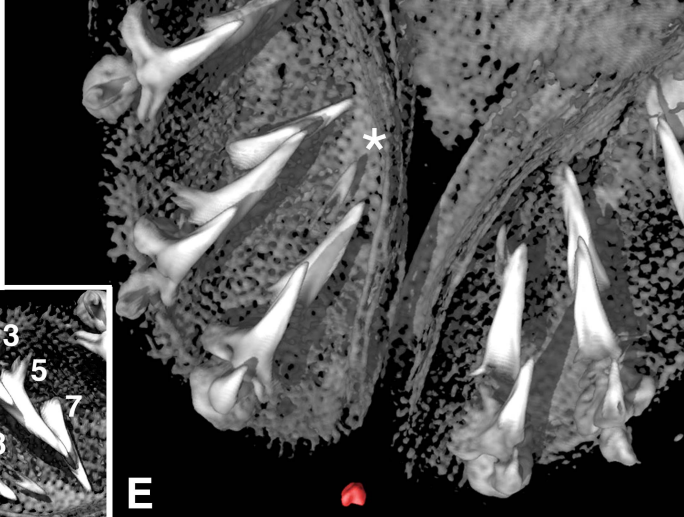
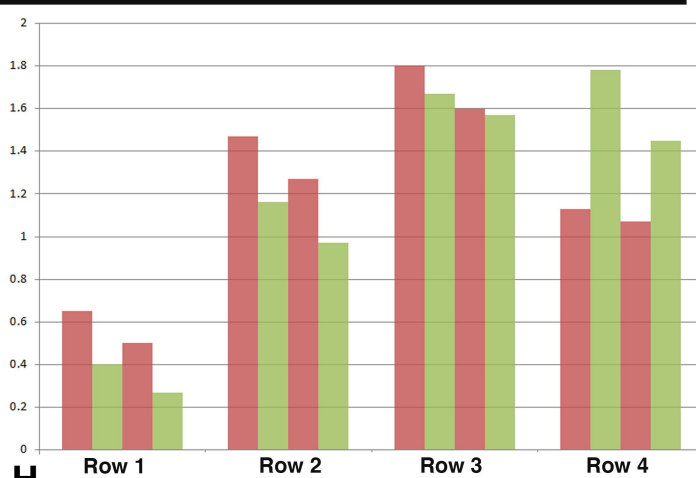
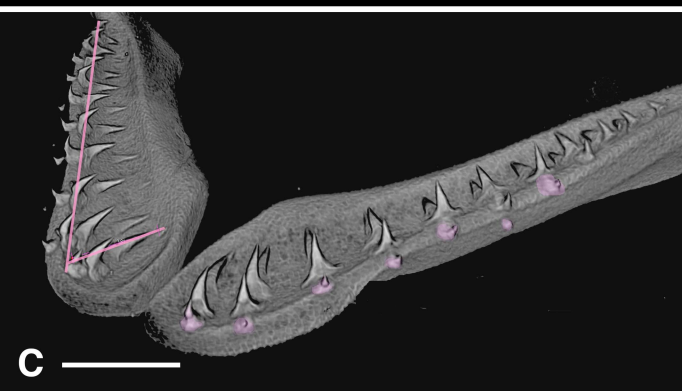
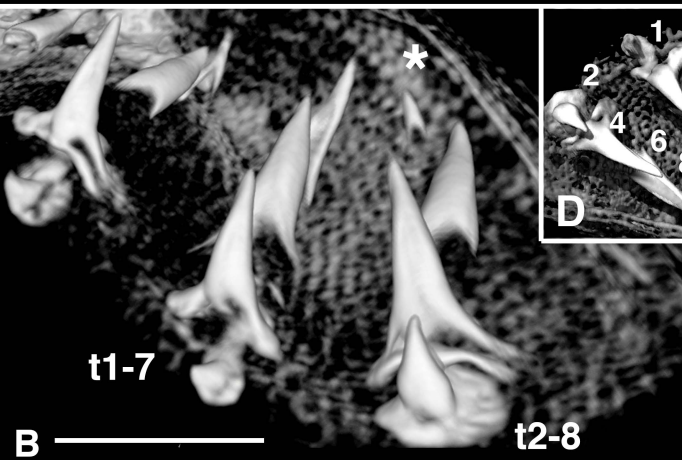
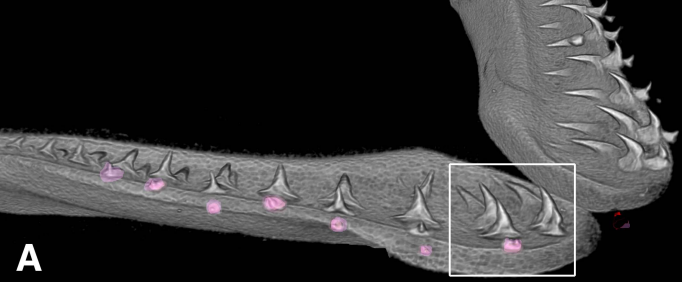
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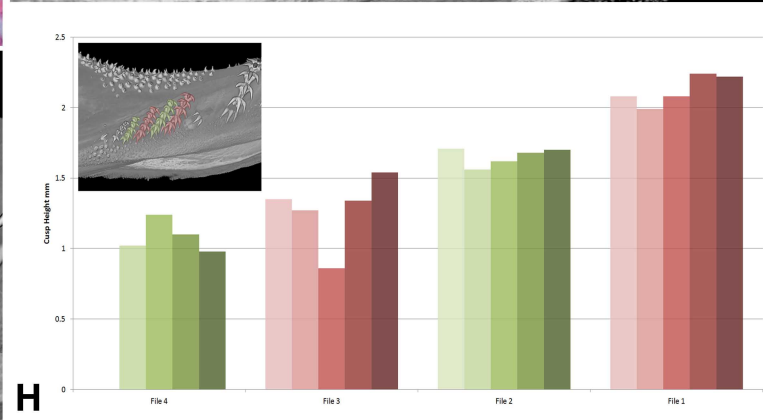
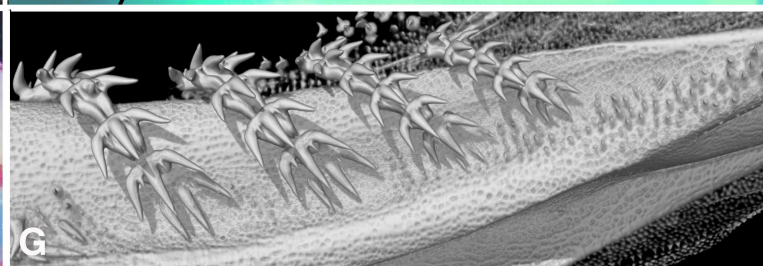
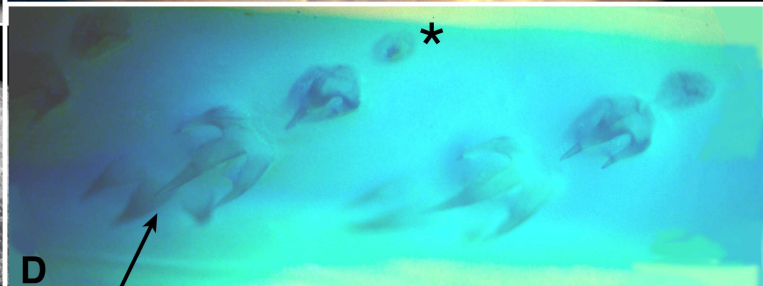
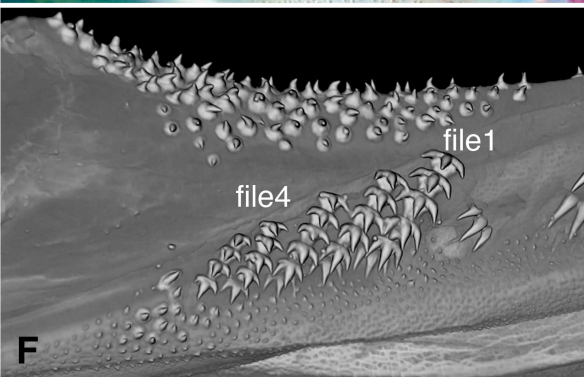
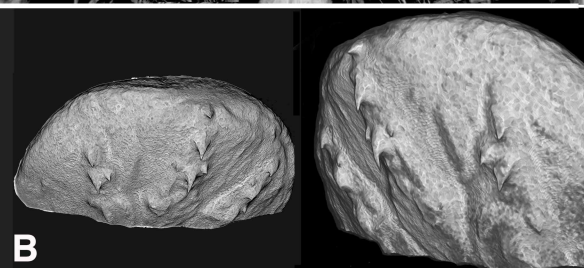
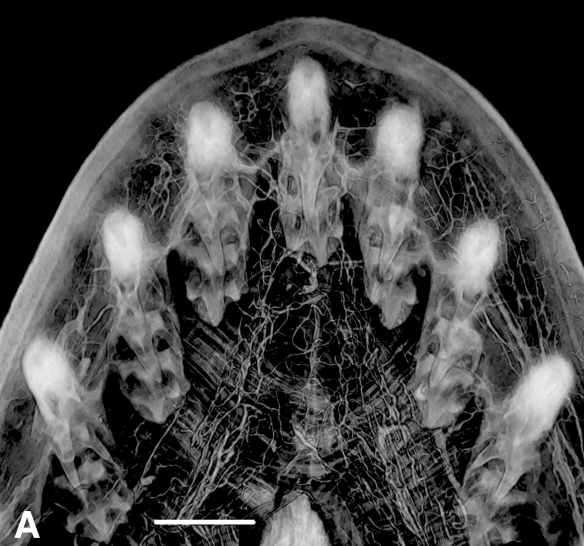
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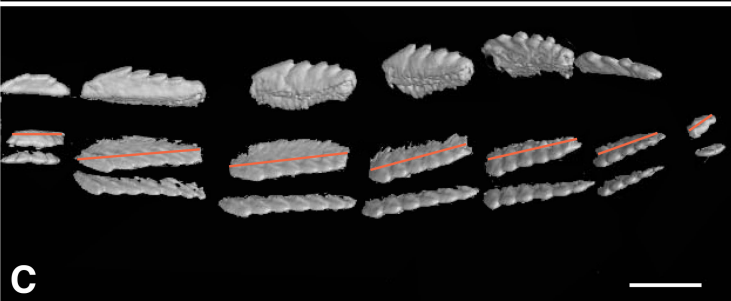


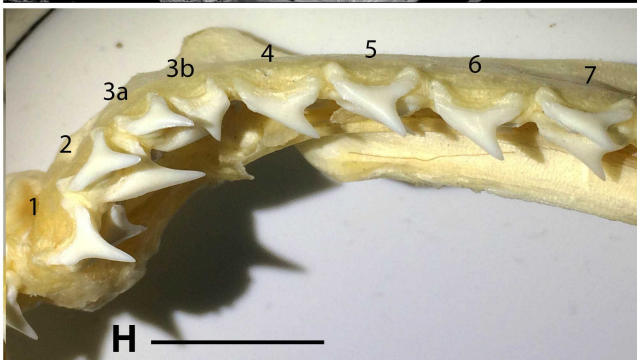
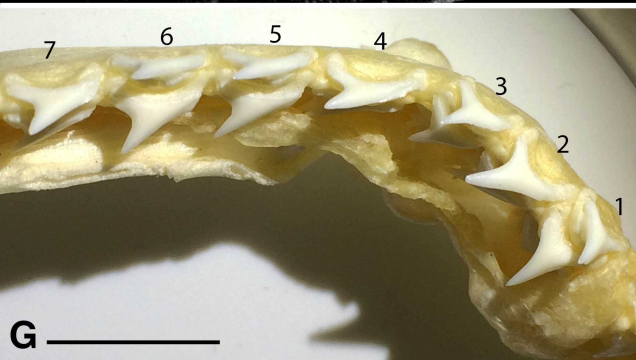
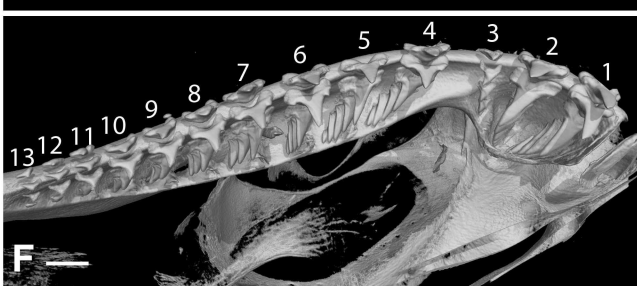
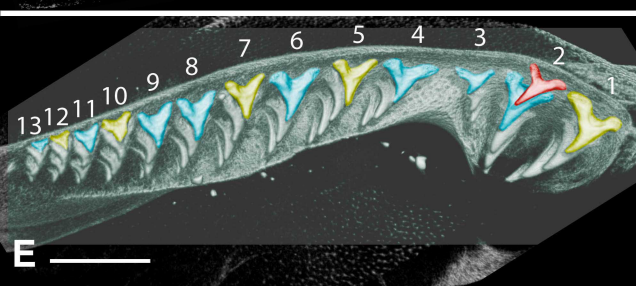
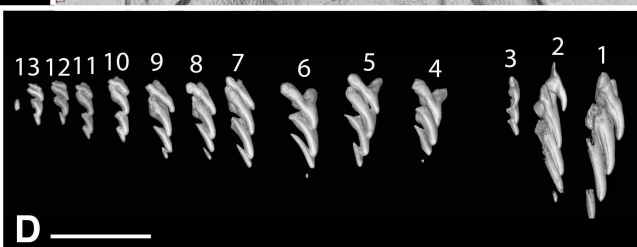
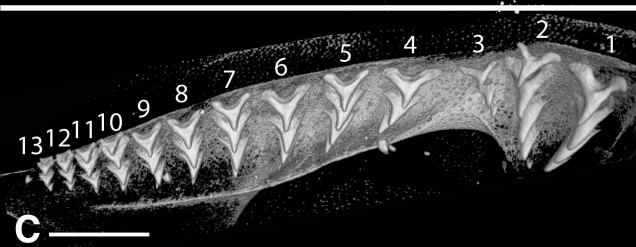
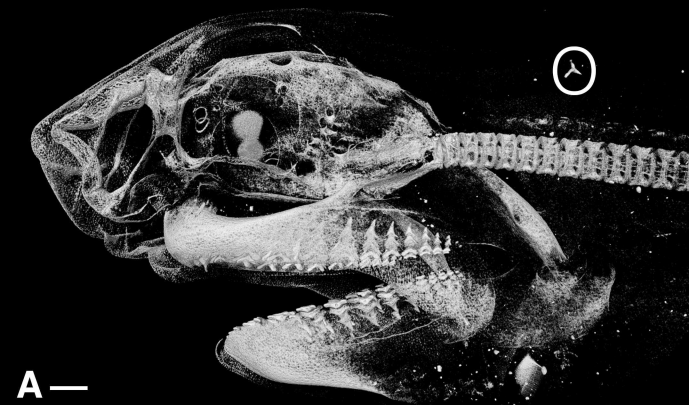


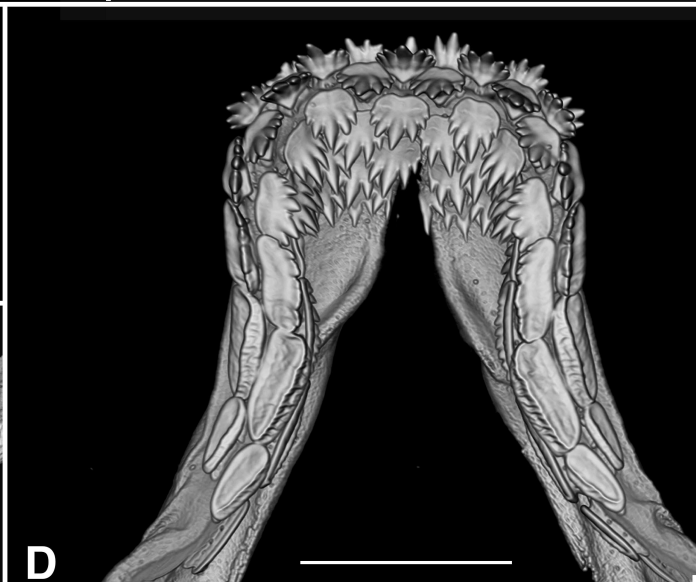
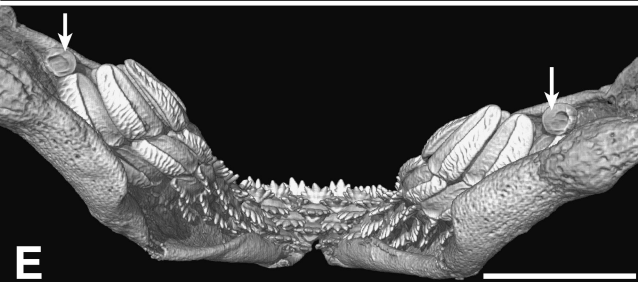
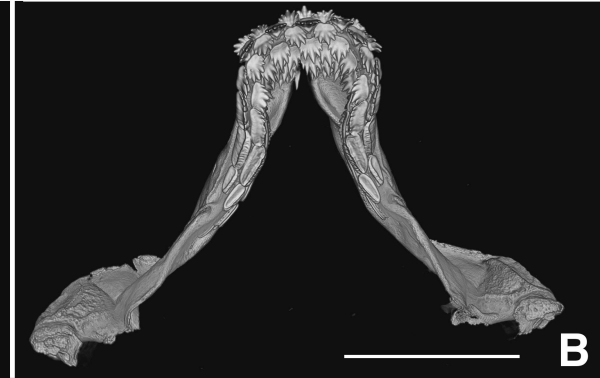


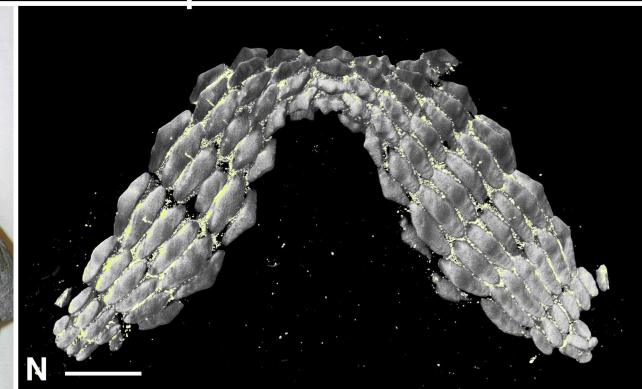
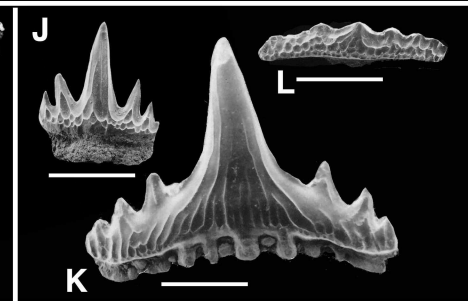
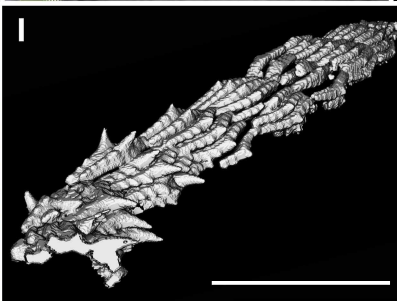
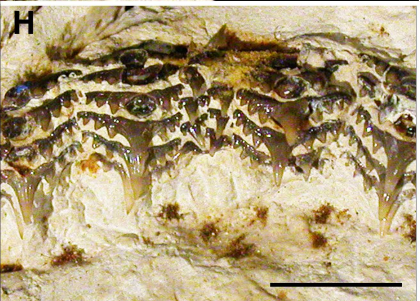
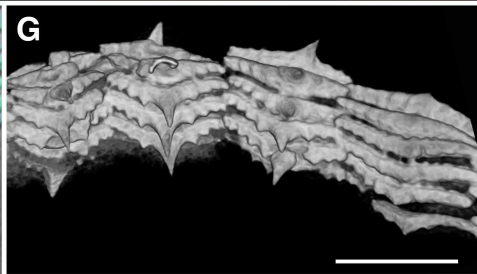
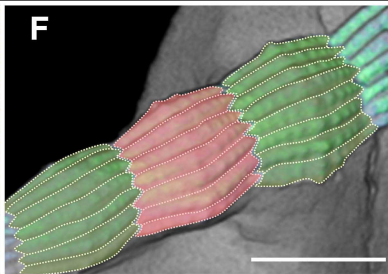
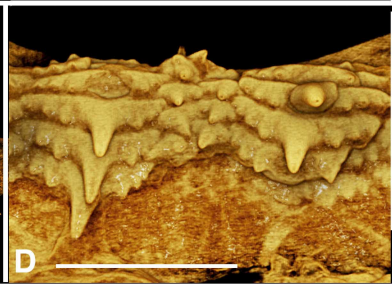
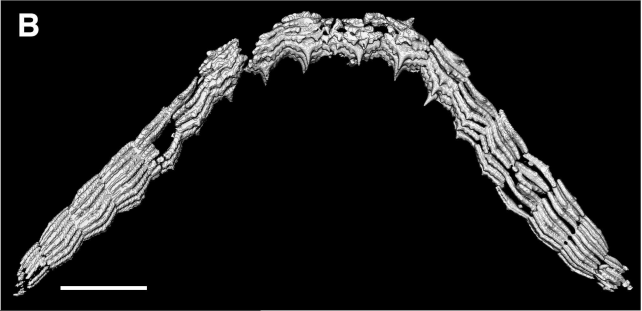
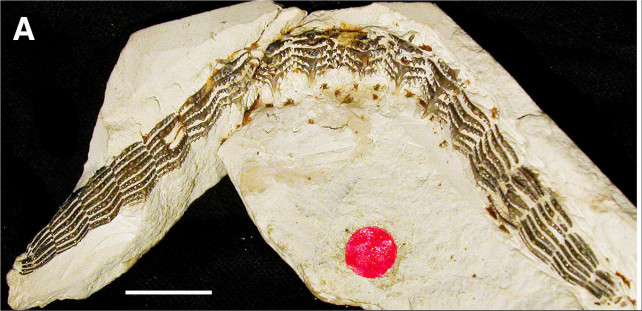
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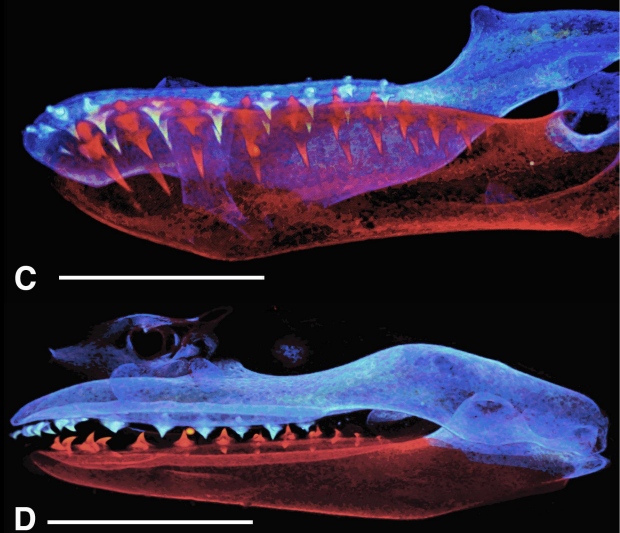
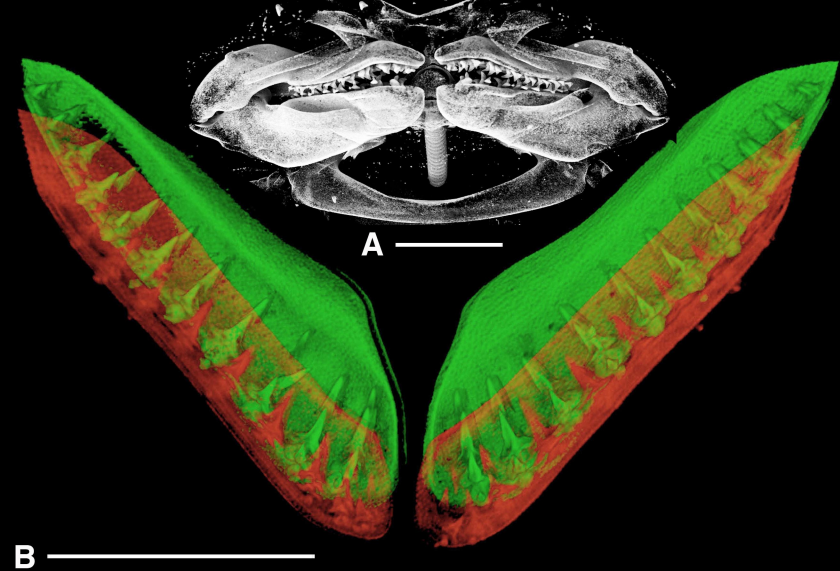


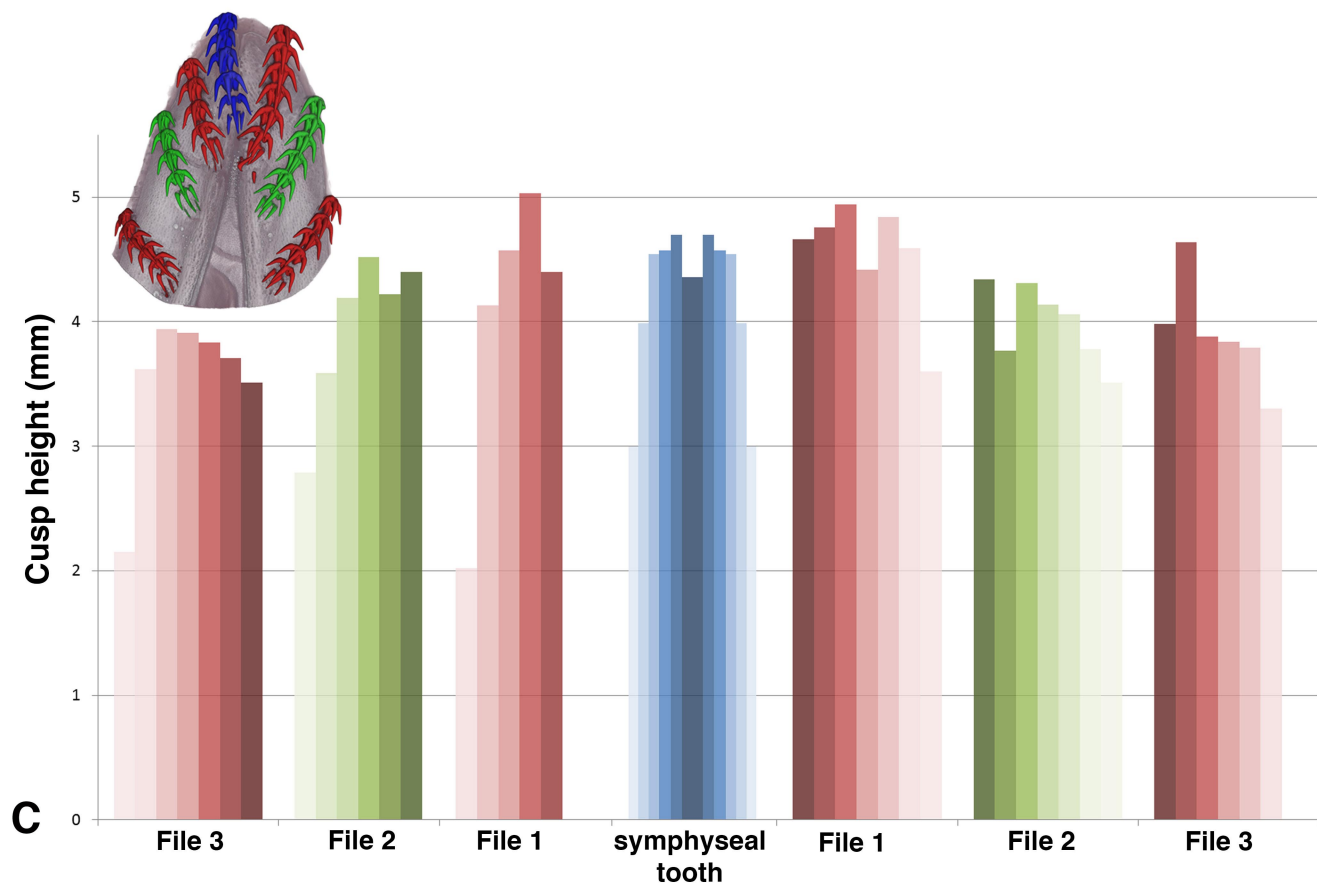
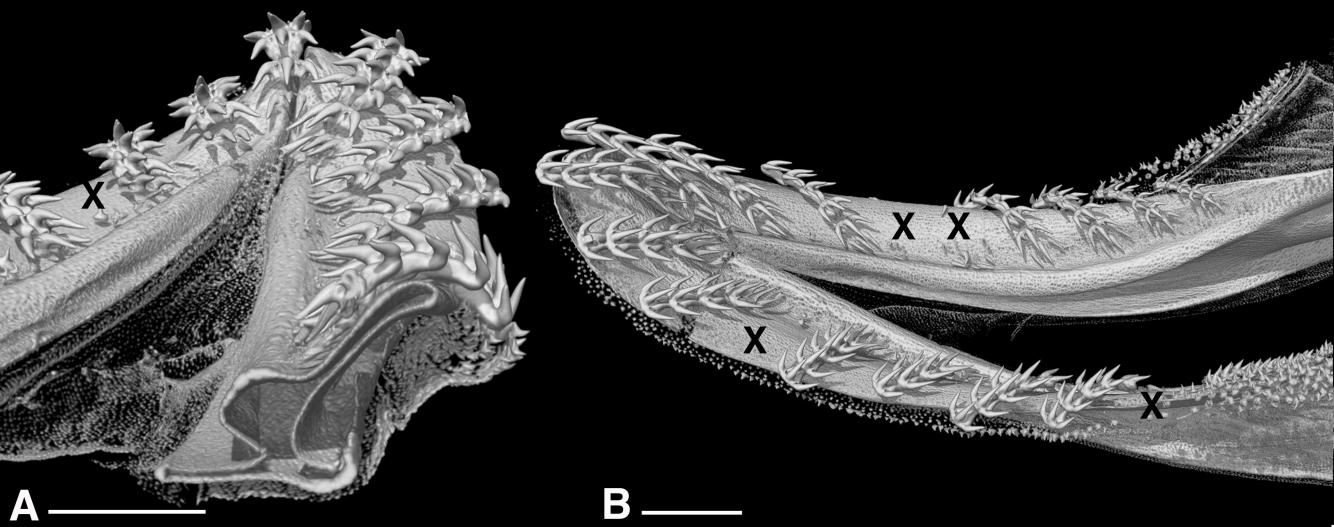


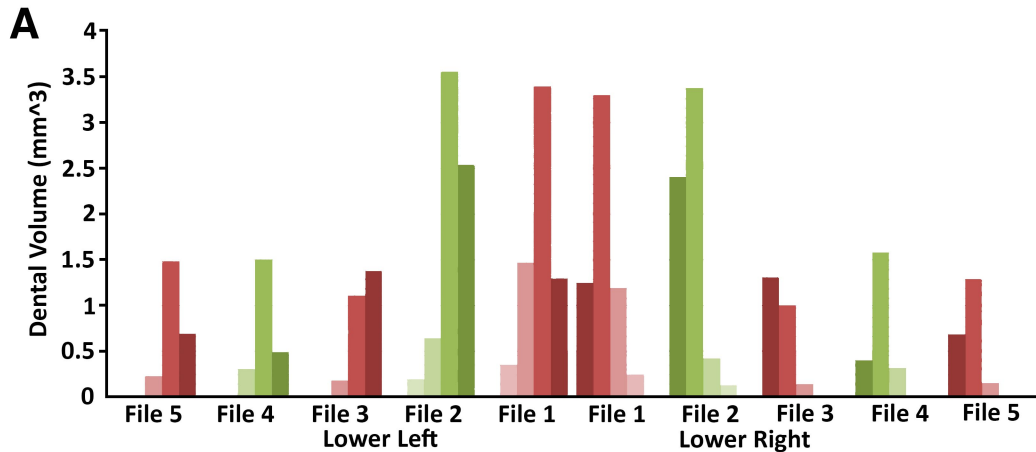




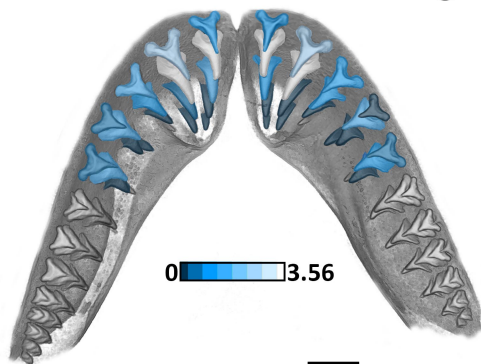








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